GaMYB1_alignment_and_sequence.txt

Exhibit 2

```
qi|7438345|pir||T09879
                                myb-related protein A - upland cotton
gi | 437327 | gb | AAA33067.1 |
                                MYB1 [Gossypium hirsutum]
gi|23476277|gb|AAN28269.1|
                                myb-like transcription factor 1 [Gossypium hirsutum]
Length = 294
Score = 602 \text{ bits } (1553), Expect = e-171
Identities = 288/294 (97%), Positives = 291/294 (98%)
Frame = +3
Query: 144
            MGRSPCCEKAHTNKGAWTKEEDORLINYIRVHGEGCWRSLPKAAGLLRCGKSCRLRWINY 323
            MGRSPCCEKAHTNKGAWTKEEDQRLINYIRVHGEGCWRSLPKAAGLLRCGKSCRLRWINY
Sbjct: 1
            MGRSPCCEKAHTNKGAWTKEEDQRLINYIRVHGEGCWRSLPKAAGLLRCGKSCRLRWINY 60
Query: 324
            LRPDLKRGNFTEEEDELIIKLHSLLGNKWSLIAGRLPGRTDNEIKNYWNTHIKRKLISRG 503
            LRPDLKRGNFTEEEDELIIKLHSLLGNKWSLIAGRLPGRTDNEIKNYWNTHIKRKLISRG
Sbjct: 61
            LRPDLKRGNFTEEEDELIIKLHSLLGNKWSLIAGRLPGRTDNEIKNYWNTHIKRKLISRG 120
Query: 504
            IDPQTHRPLNQTAITNTVTGPTELDFRNSPTSVSKSSSIKNPSLDFNYNEFHFKSHTDSL 683
            IDPQTHRPLNQTA TNTVT PTELDFRNSPTSVSKSSSIKNPSLDFNYNEF FKS+TDSL
Sbjct: 121
            IDPQTHRPLNQTANTNTVTAPTELDFRNSPTSVSKSSSIKNPSLDFNYNEFQFKSNTDSL 180
            EEPNCTASTGMTTDEEQQEQLHKKQQYGPSNGQDINLELSIGIVSADSSRVSSANSAESK 863
Query: 684
            EEPNCTAS+GMTTDEEQQEQLHKKQQYGPSNGQDINLELSIGIVSADSSRVS+ANSAESK
Sbict: 181
            EEPNCTASSGMTTDEEQQEQLHKKQQYGPSNGQDINLELSIGIVSADSSRVSNANSAESK 240
Query: 864
            PKVDNNNFQFLEQAMVAKAVCLCWQLGFGTSEICRNCQNSNSNGFYSYCRPLDS 1025
            PKVDNNNFQFLEQAMVAKAVCLCWQLGFGTSEICRNCQNSNSNGFYSYCRPLDS
Sbict: 241
            PKVDNNNFQFLEQAMVAKAVCLCWQLGFGTSEICRNCQNSNSNGFYSYCRPLDS 294
```

>Contig20 GaMYB1

GGCACGAGACAGTTTCTCTTTTTTGCTCATTTCCATCATCCCCATGTCCC CTCTCCATGTTTTTCTAAATCTCTCCATATACCTATAACACCGTTATTCT TTCTCTATTCTACCTGATTTGATTTGATTTTGTAACTGATGGGAC GATCACCTTGTTGTGAAAAGGCTCATACCAACAAGGTGCCTGGACCAAA GAGGAAGATCAACGCCTCATCAACTACATCCGTGTCCATGGTGAAGGCTG CTGGCGTTCCCTCCCAAAGCTGCTGGGCTGCTTAGATGTGGTAAGAGTT GCAGATTAAGATGGATAAACTACTTGAGGCCTGATCTTAAGAGAGGAAAT TTCACTGAAGAAGAAGATGAGCTTATCATCAAGCTTCACAGTTTACTTGG AGATAAAGAACTACTGGAACACACACATCAAAAGAAAGCTTATAAGCAGA GGAATTGATCCACAAACTCATCGTCCTCTCAATCAAACGGCCATTACCAA CACAGTCACAGGCCCCACCGAATTGGATTTCAGAAACTCGCCCACATCCG TTTCCAAATCCAGTTCCATCAAAAACCCGTCTCTGGATTTCAATTACAAT GAATTTCATTTCAAGTCCCACACAGATTCCCTTGAAGAACCCAACTGTAC AGCCAGCACTGGCATGACTACAGATGAAGAACAACAAGAACAGCTGCACA AGAAGCAGCAATACGGTCCGAGCAATGGGCAAGACATAAATTTGGAGCTG TCGATTGGGATTGTTTCAGCTGACTCATCTCGGGTATCAAGTGCCAACTC GGCCGAGTCGAAACCAAAGGTAGATAACAACAATTTCCAGTTTCTTGAAC AAGCTATGGTGGCTAAGGCGGTATGTTTGTGTTTGGCAATTAGGTTTTGGA ACAAGTGAAATTTGTAGGAACTGTCAAAATTCAAATTCAAATGGCTTCTA TAGTTATTGTAGACCCTTGGATTCATAGGGTCATCTTTTTCTTCTT CACTGTAATCAAAGCAAAATTATAAAACAAAGACTCTTTTTGATTTGTTC AATTTATAGGTTCAAAAAAAAAAAAAAAAA

GaMYB1_alignment_and_sequence.txt

>Contig20_Frame+3 GaMYB1
HETVSLFCSFPSSPCPLSMFF*ISPYTYNTVILSLFYLI*FDLIL*LMGRSPCCEKAHTN
KGAWTKEEDQRLINYIRVHGEGCWRSLPKAAGLLRCGKSCRLRWINYLRPDLKRGNFTEE
EDELIIKLHSLLGNKWSLIAGRLPGRTDNEIKNYWNTHIKRKLISRGIDPQTHRPLNQTA
ITNTVTGPTELDFRNSPTSVSKSSSIKNPSLDFNYNEFHFKSHTDSLEEPNCTASTGMTT
DEEQQEQLHKKQQYGPSNGQDINLELSIGIVSADSSRVSSANSAESKPKVDNNNFQFLEQ
AMVAKAVCLCWQLGFGTSEICRNCQNSNSNGFYSYCRPLDS*GHLFLLSFCF*EIN*FLI
IIFLPAHCNQSKIIKQRLFLICSIYRFKKKKKK

>GAMYB1 - putative from GhMYB1 alignment
MGRSPCCEKAHTNKGAWTKEEDQRLINYIRVHGEGCWRSLPKAAGLLRCGKSCRLRWINY
LRPDLKRGNFTEEEDELIIKLHSLLGNKWSLIAGRLPGRTDNEIKNYWNTHIKRKLISRG
IDPQTHRPLNQTAITNTVTGPTELDFRNSPTSVSKSSSIKNPSLDFNYNEFHFKSHTDSL
EEPNCTASTGMTTDEEQQEQLHKKQQYGPSNGQDINLELSIGIVSADSSRVSSANSAESK
PKVDNNNFQFLEQAMVAKAVCLCWQLGFGTSEICRNCQNSNSNGFYSYCRPLDS